ERRATA

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D. Gravestock, M. C. Dovey, Synthesis, 2003, 523.

We have been interested in unusual chemical shifts displayed by compounds **6a–k**. Attempts to rationalise these observations by performing chemical transformations and *ab initio* quantum chemical calculations (Gaussian 98) met with little success. It was subsequently felt that a crystal structure might reveal valuable information as to the electronic character of the compounds in question. Following further work, a crystal suitable for X-ray diffraction studies was obtained and this revealed that the original structure proposed for **6a–k** was, in fact, incorrect. The correct structure is shown below (Scheme 1) and the authors apologise for this error.

Scheme 1